

Keeping time with the human genome

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The cloning and characterization of 'clock gene' families has advanced our understanding of the molecular control of the mammalian circadian clock. We have analysed the human genome for additional relatives, and identified new candidate genes that may expand our knowledge of the molecular workings of the circadian clock. This knowledge could lead to the development of therapies for treating jet lag and sleep disorders, and add to our understanding of the genetic contribution of clock gene alterations to sleep and neuropsychiatric disorders. The human genome will also aid in the identification of output genes that ultimately control circadian behaviours.

Completion of the draft sequence of the human genome is an epic event, providing a treasure-trove of new genes for all aspects of biological investigation. For circadian clock research, 'snapshot' analysis of the draft sequence provides a glimpse of potential new members of canonical 'clock gene' families. Such candidate genes will add to the riches of the post-genomic era, ultimately leading to a wider understanding of the molecular control of the circadian clock.

The master circadian clock in mammals, in the suprachiasmatic nuclei (SCN) of the anterior hypothalamus, drives daily variations in many physiological and behavioural processes, such as the sleep-wake rhythm and daily variations in body temperature, hormone levels, cognition and memory¹. Dawn and dusk coordinate or entrain the circadian clock through neural pathways connecting the retina to the SCN, so that the master clock and its output rhythms do not drift from 24 h, but remain aligned with the solar day. Transient disruption of circadian timing following transmeridian flights leads to jet lag, and chronic alterations of the central clock mechanism in shift workers (around 25% of the working population) may contribute to poor health and sleep disorders. Finally, specific rhythm defects may be involved in neuropsychiatric illnesses.

The fruitfly *Drosophila melanogaster* has long been the animal model of choice for genetic analysis of circadian rhythms. In recent years, with the help of forward and reverse genetics, homology-based approaches and protein-interaction screens, homologues of most of the genes involved in the fly clock have been cloned in mammals² (Fig. 1, Table 1). Although there are compelling similarities between the general clock mechanisms of *Drosophila* and mice, gene duplication has led to reassignment of specific functions among structurally homologous components.

Analysis of the fly circadian clock appears to be close to genetic saturation, indicating that most of the cardinal genes may have been identified³. It is less clear whether the whole complement of clock genes has been identified in mammals, because expansion of gene families makes it likely that additional family members may await discovery. We searched the draft sequence of the human genome for new clock genes (see Supplementary Information for Methods).

A few words of caution are in order concerning the sequence analysis. Although we have identified new clock gene candidates, we have no information regarding their expression patterns or function. We detected all of the previously identified clock loci in mammals, confirming the near completeness of the human genome sequence data. This does not rule out the possibility, however, of further clock candidates surfacing from advanced searches of the finished genome sequence.

Genes involved in the core clock mechanism

The core clock mechanism of mammals is composed of interacting positive and negative transcriptional/translational feedback loops⁴ (Fig. 1). The pivotal components of this mechanism are two basic helix-loop-helix (bHLH)/PAS-containing transcription factors, CLOCK and BMAL1, which are assumed to pair through their protein-interactive PAS domains. PAS domains are found in a diverse family of proteins (PER, ARNT and SIM are the founder members) and are a common feature of transcription factors that are essential for the clock machinery of fungi, insects and mammals⁵.

CLOCK-BMAL1 heterodimers drive the rhythmic transcription of three *Period* genes (*mPer1–mPer3* in the mouse) and two

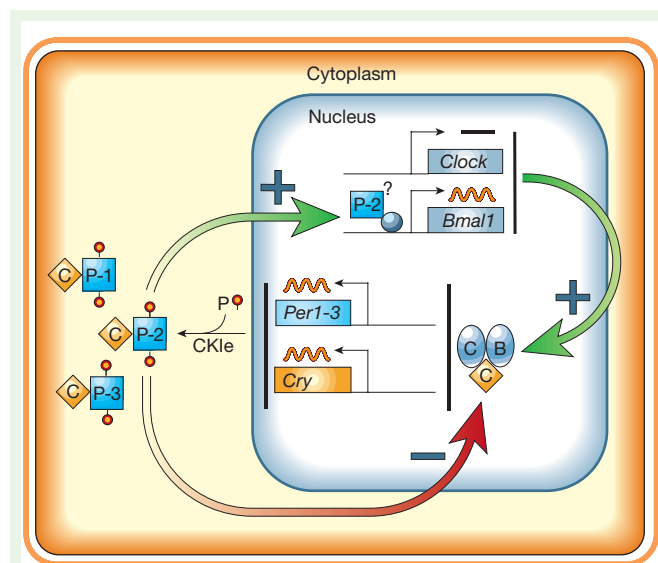


Figure 1 Model of circadian clockwork within the SCN. The clockwork is comprised of interacting positive (green) and negative (red) feedback loops. CLOCK (oval with C) and BMAL1 (oval with B) heterodimers activate (+) rhythmic transcription of *Cry* and *Per* genes. The CRY (C) and PER (P1–P3) proteins form complexes that are important for nuclear translocation of the PER proteins. The phosphorylation (P) of the PER proteins by CKIε may also regulate their cellular location and stability. PER2 may positively regulate transcription of *Bmal1* by acting as a co-activator. The nuclear-localized mCRY proteins interact directly with CLOCK and BMAL1 to negatively regulate (–) transcription mediated by CLOCK–BMAL1. PER1 may communicate light information to the molecular loops, as its RNA can be induced by light at dawn and dusk. PER3 may contribute to transducing the oscillation to output systems.

Cryptochrome genes (*mCry1* and *mCry2*). As the mPER and mCRY proteins are translated, they form PER–CRY complexes that are translocated to the nucleus. There the mCRY proteins act as negative regulators by directly interacting with CLOCK or BMAL1, or both, to inhibit transcription, forming a negative feedback loop. At the same time, mPER2 contributes to the transcription of *Bmal1*, which is rhythmically expressed with a peak phase opposite to that of *mPer/mCry*, forming a positive feedback loop. The push–pull action of the positive and negative feedback loops perpetuates the self-sustaining nature of the circadian clock.

Analysis of the draft sequence identified human *hPER1*, *hPER2*, *hPER3*, *hCRY1* and *hCRY2*. Moreover, an additional single-copy gene was identified on chromosome 7 with high sequence similarity to the *Per* family (Table 1). A fragment of genomic sequence is available, but no expressed sequence tag (EST) matches were obtained. Nevertheless, the putative translation of this fragment encodes 231 residues that encompass most of the PAS region and can be compared with those of other family members (see Supplementary Information Figs A and B). If '*hPER4*' can be confirmed as an expressed sequence rather than a pseudogene, the human genome project will have yielded a potentially exciting finding, because such a fourth *PER* will contribute to our understanding of the core clock mechanism. Even if it is not expressed, identification of an *hPER4* locus may help us to decipher the evolution of the *Per* gene family.

Our analysis identified *hBMAL1* (*hMOP3*) and *hCLOCK*, as well as two previously described members of this transcription factor family, *hMOP4* and *hMOP9* (also called *CLIF*)^{6–8}, which encode proteins closely related to hCLOCK and hBMAL1, respectively (Table 1; see Supplementary Information Figs C and D). We did not find any new genes related to these transcription factors. Because MOP4–BMAL1 and CLOCK–MOP9/CLIF heterodimers can activate transcription from E-boxes (CACGTG) *in vitro*, MOP4 and MOP9/CLIF have the potential to contribute to transcriptional regulation within the core clock mechanism. MOP9/CLIF is in fact expressed in the SCN^{7,8}.

Kinases important for clock function

Phosphorylation and proteolysis of clock proteins can determine their cellular location and stability, and are key ingredients for building time delays into the 24-h molecular mechanism³. In *Drosophila*, DOUBLETIME, which is closely related to mammalian casein kinase I epsilon (CKIε), phosphorylates PER, thereby influencing PER turnover³. Studies of the *tau* mutation in Syrian hamsters (a spontaneous, semidominant mutation leading to marked shortening of the circadian period) reveal that it encodes a missense mutation within CKIε that renders the mutant enzyme deficient in its ability to phosphorylate the mPER proteins⁹.

Casein kinase I delta (CKIδ) is highly homologous to CKIε (76% identical at the amino-acid level) and efficiently binds and phosphorylates mPER1 *in vitro*¹⁰. Our database search revealed another human casein kinase gene, '*hCKIε2*'. This gene is detected in three partially overlapping ESTs that could encode a carboxy-terminal fragment highly homologous to hCKIε (91% identical over 73 residues, with two of the substitutions in or near putative autophosphorylation sites; see Supplementary Information Figs E and F). Three other novel *hCK1* genes were found, one similar to chicken *CK1γ1* (which we name *hCK1γ1*), and two related to *hCK1α1*, which we have called *hCK1α2* and *hCK1α3* (see Supplementary Information Figs E and F). Functional analysis of CKIε2 will certainly add to the complexity inherent in the phosphorylation events that are important for the clockwork.

The timeless mystery

The *timeless* (*tim*) gene is essential for circadian function in *Drosophila*³. Putative homologues of *Drosophila tim* have been identified in both mice and humans (*mTim* and *hTIM*, respectively),

but placing the homologue within the murine molecular mechanism has proved difficult². Database searches of the completed *Drosophila* genome show that mammalian TIM is not the true orthologue of *Drosophila* TIM, but is the likely orthologue of a newly described fly gene, *timeout* (also called *tim-2*)^{11,12}. As the core clock functions of TIM in the fly have been usurped by other clock-relevant genes in the mouse², there is no obvious requirement for a mammalian equivalent. Our analysis of the human genome sequence did not reveal a second *Timeless* homologue, consistent with this idea.

Analysis of the completed genome of the worm *Caenorhabditis elegans* provides additional insight into the *timeless* mystery. These animals are behavioural outliers, because no circadian rhythms have been reported in the worm. We might reasonably predict that some of the canonical clock genes would be absent from *C. elegans*. In fact, our searches of the worm databases did not detect any genes with similarity to cryptochromes. Significant hits were obtained for both *clock* and *bmal1* to the same *C. elegans* gene *aha-1* (an *hARNT* homologue; Table 1), with the similarity far greater for *bmal1*. The *C. elegans* gene *lin-42*, which is required for postembryonic development, shows some similarity in its PAS region to the *Drosophila* and mammalian PERs¹³.

Table 1 Clock genes in metazoans

Fruittfly (<i>Drosophila melanogaster</i>)	Worm (<i>Caenorhabditis elegans</i>)	Mouse (<i>Mus musculus</i>)	Human (<i>Homo sapiens</i>)
Period			
–	–	<i>mPer1</i> [11B] SWALL: O35973	<i>hPER1</i> [17p13.1] SWALL: O15534
<i>per</i> SWALL: P07663	<i>lin-42</i> SWALL: P91313	<i>mPer2</i> SWALL: O54943	<i>hPER2</i> [2] SWALL: O15055
–	–	<i>mPer3</i> SWALL: O70361	<i>hPER3</i> [1] SWALL: Q9UGU8
–	–	–	<i>hPER4</i> [7] gb: AC027390.3
Timeless			
<i>tim</i> SWALL: P49021	–	–	–
<i>timeout/tim2</i> SWALL: Q9NG27	<i>tim-1</i> SWALL: Q9XW65	<i>mTim</i> * [10:18.0] SWALL: Q9Z0E7	<i>hTIM</i> [12q12] SWALL: O94802
Clock/Bmal1			
<i>Clock/Jerk</i> SWALL: O61735	<i>aha-1</i> SWALL: Q18141	<i>mClk</i> [5:43.0] SWALL: O08785 <i>mNPAS2</i> [1.20] SWALL: P97460	<i>hCLK</i> [4q12] SWALL: O15516 <i>hMOP4</i> [2p11.2] SWALL: Q99742
<i>cycle</i> SWALL: O61734	<i>aha-1</i> SWALL: Q18141	<i>mBmal1</i> (<i>mMOP3</i>) [7.52] SWALL: Q9WTL8 SWALL: O88295	<i>hBMAL1</i> (<i>hMOP3</i>) [11p15] SWALL: O00327
–	–	–	<i>hMOP9</i> (<i>CLIF</i>) [12p12.2] SWALL: Q9NYQ5
Casein Kinase 1ε			
<i>doubletime</i> * SWALL: O76324	<i>CK1δ</i> SWALL: Q20471	<i>tau</i> (<i>mCK1ε</i>) [15:E1] SWALL: Q9JMK2	<i>hCK1ε</i> [22q13.1] SWALL: P49674
–	–	–	<i>hCK1ε2</i> aw673521 aa934662 aa001679
Cryptochrome			
<i>cry</i> SWALL: O77059	–	<i>mCry1</i> [10C] SWALL: P97784	<i>hCRY1</i> [12q23] SWALL: Q16526
–	–	<i>mCry2</i> [2E] SWALL: Q9R194	<i>hCRY2</i> [11] SWALL: O75148

The results are from searches of the draft human genome database with members of the canonical clock genes from fly and mouse. Genes on the same row are the most similar between species: for example, *dper* is most similar in sequence to *mPer2*, and worm *CK1δ* is the closest to fly and mammalian *CKIε*. Dashes indicate that there is no similar sequence in the corresponding database. No similarities to either *hPER4* or *hCKIε2* were found in mouse genomic or expressed sequence tag databases. For the human genome, chromosome positions (if known) are given in square brackets, and novel, closely related family members discovered in this search are given in bold. Underlined genes, when mutated, give clock phenotypes. SWALL, SwissProt accession numbers; gb, GenBank accession numbers. See Supplementary Information for detailed Methods.

* Loci with lethal mutations associated.

Most notably, the worm has only one *timeless* gene, *tim-1*, which is highly homologous to *Drosophila timeout/tim-2* and mammalian *Tim*, but substantially removed from the clock-relevant *timeless* in the fly¹³. This might suggest that the closely related worm *tim-1*, mouse/human *Tim* and fly *timeout/tim-2* genes are descendants of an ancestral *timeless* gene that duplicated in the arthropod lineage after the split with nematodes and vertebrates¹². The clock-relevant *timeless* duplication could then have rapidly diverged to take on a dedicated role in the circadian machinery of insects.

Impact of the human genome on chronobiology

This snapshot analysis of the draft sequence of the human genome provides candidate sequences that could increase the number of clock genes in mammals. Aided by further genomic and post-genomic analyses, this could provide new opportunities for pharmacological manipulation of the human clockwork with 'chronotherapies' targeted at improving the treatments for those who work shifts or suffer from jet lag, and those with clock-related sleep or psychiatric problems.

Recently, an autosomal dominant sleep disorder characterized by debilitating early sleep onset and offset was shown to involve a missense mutation in the CKIε binding region of *hPER2*; the mutation leads to decreased phosphorylation of *hPER2* by CKIε *in vitro*¹⁴. Linkage analysis identified a candidate locus in the affected family, and the chromosomal location of known clock genes from the human genome provided a short-cut in the search. Identified polymorphisms within human clock genes may aid genetic analysis of other rhythm-related traits, such as the early riser 'lark' and the late riser 'owl' phenotypes¹⁵.

The completed human genome sequence will also have a significant impact on the study of circadian output systems. One of the most profound discoveries of the mammalian clock gene 'era' has been the finding that clock genes in mice and humans are widely expressed throughout the body, where they seem to participate in local circadian oscillator function². These peripheral oscillators are dampened, and require cyclic input from the master clock in the SCN for sustained rhythmicity. Local clock-controlled genes in peripheral organs would provide the flexibility needed to control the circadian rhythms present in these tissues.

Using DNA chip technology, expression profiling of rhythmic gene expression in the SCN and in peripheral tissues will identify candidate output genes and their temporal contribution to local physiology. The completed human genome sequence will then play its part in the identification of these output genes and accelerate fulfilment of one of the great promises of circadian rhythm research: providing a complete understanding of the cellular and molecular events connecting clock genes to circadian behaviour.

1. Klein, D. C., Moore, R. Y. & Reppert, S. M. (eds) *Suprachiasmatic Nucleus: The Mind's Clock* (Oxford Univ. Press, New York, 1991).
2. Reppert, S. M. & Weaver, D. R. Comparing clockworks: mouse versus fly. *J. Biol. Rhythms* **15**, 357–364 (2000).
3. Young, M. W. Circadian rhythms. Marking time for a kingdom. *Science* **288**, 451–453 (2000).
4. Shearman, L. P. *et al.* Interacting molecular loops in the mammalian circadian clock. *Science* **288**, 1013–1019 (2000).
5. Dunlap, J. C. Molecular bases for circadian clocks. *Cell* **96**, 271–290 (1999).
6. Hogenesch, J. B., Gu, Y. Z., Jain, S. & Bradfield, C. A. The basic-helix-loop-helix-PAS orphan MOP3 forms transcriptionally active complexes with circadian and hypoxia factors. *Proc. Natl Acad. Sci. USA* **95**, 5474–5479 (1998).
7. Hogenesch, J. B. *et al.* The basic helix-loop-helix-PAS protein MOP9 is a brain-specific heterodimeric partner of circadian and hypoxia factors. *J. Neurosci.* **20**, RC83 (2000).
8. Maemura, K. *et al.* CLIF, a novel cycle like factor, regulates the circadian oscillation of plasminogen activator inhibitor-1 gene expression. *J. Biol. Chem.* **275**, 36847–36851 (2000).
9. Lowrey, P. L. *et al.* Positional syntenic cloning and functional characterization of the mammalian circadian mutation *tau*. *Science* **288**, 483–491 (2000).
10. Vielhaber, E., Eide, E., Rivers, A., Gao, Z. H. & Virshup, D. M. Nuclear entry of the circadian regulator mPER1 is controlled by mammalian casein kinase 1ε. *Mol. Cell. Biol.* **10**, 4888–4899 (2000).
11. Gotter, A. L. *et al.* A time-less function for mouse *Timeless*. *Nature Neurosci.* **3**, 755–756 (2000).
12. Benna, C. *et al.* A second *timeless* gene in *Drosophila* shares greater sequence similarity with mammalian *tim*. *Curr. Biol.* **10**, R513 (2000).
13. Jeon, M., Gardner, H. F., Miller, E. A., Deshler, J. & Rougvie, A. E. Similarity of the *C. elegans* developmental timing protein LIN-42 to circadian rhythm proteins. *Science* **286**, 1141–1146 (1999).
14. Toh, K. L. *et al.* An *hPer2* phosphorylation site mutation in familial advanced sleep-phase syndrome. *Science*, 11 January 2001 (10.1126/science.1057499).
15. Katzenberg, D. *et al.* A CLOCK polymorphism associated with human diurnal preference. *Sleep* **21**, 569–576 (1998).

Supplementary information is available on Nature's World-Wide Web site (<http://www.nature.com>) or as paper copy from the London editorial office of *Nature*.

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